

1 Method for the surface treatment of a titanium dioxide pigment

2 CROSS REFERENCE TO RELATED APPLICATIONS

3 This application claims priority pursuant to Title 35, United States Code,
4 Section 119(a)- (d) or (f), or 365(b) to the German Patent Application Number DE 102 36
5 366 filed 8 August, 2002, where the above named application is hereby incorporated
6 herein by reference in its entirety including incorporated material.

7 FIELD OF THE INVENTION

8 The field of the invention is the field of methods for the surface treatment of
9 titanium dioxide pigments to produce a titanium dioxide pigment with high greying
10 resistance and high hiding power, and its use in the production of decorative laminating
11 paper.

12 BACKGROUND OF THE INVENTION

13 Decorative laminating paper is an element of a decorative thermoset surface, which is used
14 with preference for finishing furniture surfaces and for laminate floorings. Laminate is the
15 term used for materials in which wood and paper, for example, are pressed with resin. The
16 use of special synthetic resins results in extraordinarily high resistance of the laminates to
17 scratching, impact, chemicals and heat.

18 The use of special-purpose papers (decorative laminating papers) permits the production
19 of decorative surfaces, where the decorative laminating paper serves not only as facing
20 paper for unattractive wood material surfaces, but also as a carrier for the synthetic resin.
21 The requirements imposed on decorative laminating paper include, for example, hiding
22 power (opacity), light-fastness (greying resistance), colour-fastness, wet strength,
23 impregnability and printability.

24 In principle, a pigment based on titanium dioxide is eminently suitable for achieving the

1 necessary opacity of the decorative laminating paper. As a rule, a titanium dioxide pigment
2 or a titanium dioxide pigment suspension is mixed with a fibre suspension during paper
3 production. The interactions between the individual components (fibres, pigment, water)
4 contribute to formation of the paper sheet and determine the retention of the pigment. The
.5 term retention refers to the retention of all inorganic substances in the paper during production.
6 In addition to the pigment and fibres used as feedstock, auxiliaries and additives are
7 generally also used. These may affect the mechanisms of interaction between the fibres,
8 the pigment and the water.

9 A number of titanium dioxide pigments exists for applications in decorative laminating
10 paper. Alongside the most important properties, such as retention and opacity (hiding
11 power), the greying resistance also plays a decisive role.

12 It is generally known that titanium dioxide is photochemically active. When exposed to UV
13 radiation in the presence of moisture, decorative laminating paper pigmented with titanium
14 dioxide displays increasing greying. To avoid this problem, the surface of the pigments is
15 treated with various substances, for instance with Al_2O_3 aquate and a colourless metal
16 phosphate (US 3 926 660), with zinc phosphate (US 5 114 486), with cerium phosphate
17 and aluminium phosphate (GB 2 042 573), or only with aluminium phosphate (EP 0 753
18 546 A2). DE 15 92 873 describes a method for improving the light-fastness of pigments,
19 where calcination at 600 °C is performed following coating with magnesium silicate.

20 TiO_2 pigments with improved retention properties, having a cores coated with consecutive
21 layers of aluminium oxide phosphate, aluminium oxide and magnesium oxide, are
22 presented in EP 0 713 904 B1, US 5,665,366 and US 5,942,281.

23 A TiO_2 pigment, having a core coated with consecutive layers of zirconium hydroxide or
24 oxyhydroxide, titanium hydroxide or oxyhydroxide, and co-precipitated phosphate and silica, and
25 finally a layer of aluminum oxyhydroxide and magnesium oxide, is shown to protect the organic
26 pigment binder from light and to decrease the loss of gloss in paint in US 6,200,375.

1 All of the preceding publications, patents, and patent applications are hereby included in
2 their entirety in this application.

OBJECTS OF THE INVENTION

-4 It is an object of the invention to produce a method capable of producing pigments
5 with high hiding power and simultaneously high greying resistance.

6 It is an object of the invention to produce a method capable of producing pigments
7 with high hiding power and simultaneously high greying resistance for use in decorative
8 laminating papers.

SUMMARY OF THE INVENTION

10 The steps of the most preferred method of the invention are:

- 11 a) Addition of a phosphorus compound to an aqueous suspension of titanium dioxide base
12 material;
13 b) Addition of a titanium compound;
14 c) Addition of an aluminium compound;
15 d) Adjustment of the pH value of the suspension to a pH value of 8 to 10, preferably 8.5
16 to 9.5;
17 e) Addition of a magnesium compound;
18 f) Stabilisation of the pH value of the suspension in the range from 8 to 10, preferably 8.5
19 to 9.5; and
20 g) Separation of the TiO_2 pigment by filtration, followed by washing, drying and milling of
21 the pigment.

DETAILED DESCRIPTION OF THE INVENTION

24 Using pigments with improved greying resistance produced by prior art methods generally
25 reduces retention and/or opacity in laminates. Prior art improvements in opacity are

1 accompanied by a deterioration in greying resistance.

2 Therefore, the object of the present invention is to offer a method capable of producing
3 pigments with high hiding power and simultaneously high greying resistance for use in
4 decorative laminating papers.

5 The object is solved by subjecting a titanium dioxide pigment to a surface treatment
6 process characterised by the following steps:

- 7 a) Preparation of an aqueous suspension of titanium dioxide base material,
- 8 b) Addition of a phosphorus compound,
- 9 c) Addition of a titanium compound,
- 10 d) Addition of an aluminium compound,
- 11 e) Adjustment of the pH value of the suspension to a pH value of 8 to 10, preferably 8.5
- 12 to 9.5,
- 13 f) Addition of a magnesium compound,
- 14 g) Stabilisation of the pH value of the suspension in the range from 8 to 10, preferably 8.5
- 15 to 9.5,
- 16 h) Separation of the TiO_2 pigment by filtration, followed by washing, drying and milling of
- 17 the pigment.

18 Other advantageous versions of the method are described in the dependent claims of the
19 invention.

20 The object of the invention is, therefore, a method for the surface treatment of titanium
21 dioxide pigments that results in pigments with high hiding power and high retention with
22 simultaneously high greying resistance, as well as a pigment with these properties and the
23 use of this pigment in the production of decorative laminating paper.

24 Surprisingly, it was found that the addition of a titanium compound during the inorganic
25 surface treatment of the pigment with phosphorus and aluminium compounds is capable

1 of achieving both high opacity and very good greying resistance. No zirconium compounds
2 were added to the suspension to achieve this result.

3 The surface treatment process is based on TiO_2 base material, preferably produced by the
4 chloride process. The term TiO_2 base material refers to the raw TiO_2 pigment prior to post-
5 treatment. The base material can first be milled, for example in a wet-milling process. A
6 dispersant is preferably added during wet-milling. The milled base material is used to
7 prepare an aqueous suspension. This suspension can be basic or acidic and is preferably
8 basic with a pH value of 9 to 11. The method is performed at a temperature of less than
9 70 °C, preferably at 55 to 65 °C.

10 First, a phosphorus compound is added to the suspension in a quantity of 0.4 to 6.0% by
11 weight, preferably 1.0 to 4.0% by weight, calculated as P_2O_5 , referred to TiO_2 base
12 material. Particularly good results are obtained with P_2O_5 contents of 1.6 to 2.8% by weight,
13 referred to the base material. Other suitable phosphorus compounds are preferably
14 inorganic phosphorus compounds, such as alkali phosphates, ammonium phosphate,
15 polyphosphates, phosphoric acid or, where appropriate, mixtures of these compounds.
16 Other phosphorus compounds can, however, also be used.

17 Second, a titanium compound is added, e.g. titanyl sulphate, titanyl chloride or another
18 hydrolysable titanium compound, or mixtures of these compounds. The quantity of titanium
19 compound added is 0.1 to 3.0% by weight, preferably 0.1 to 1.5% by weight, and
20 particularly 0.1 to 1.0% by weight, calculated as Ti O_2 referred to TiO_2 base material in
21 the suspension.

22 Third, an aluminium compound of acidic or basic character is preferably subsequently
23 added to the suspension. Particularly suitable as an acidic aluminium compound is
24 aluminium sulphate, although this is not to be taken as a restriction. Suitable alkaline
25 aluminium compounds include sodium aluminate, alkaline aluminium chloride, alkaline
26 aluminium nitrate or other alkaline aluminium salts, or mixtures of these compounds.

- 1 The suspension will customarily be stirred for about 30 minutes following each addition, in
2 order to achieve homogenisation. It is, however, also possible to add the titanium
3 compound and the aluminium compound simultaneously.
- 4 In a preferred embodiment of the method, an acid or a base, or a second aluminium
5 compound, is added in parallel with the aluminium compound, in order to maintain a
6 constant pH value in the range from 2 to 10, preferably in the range from 4 to 9 and
7 particularly in the range from 6 to 8. In a particularly advantageous version of the method,
8 the pH value is controlled by the balanced, parallel addition of sodium aluminate and HCl.
9 A further procedure consists in keeping the pH value constant by means of the controlled
10 addition of aluminium sulphate and sodium aluminate.
- 11 The suspension is subsequently adjusted to a pH value of 8 to 10, preferably 8.5 to 9.5.
12 A person skilled in the art adjusts the pH value in the customary manner with the help of
13 appropriate acidic or alkaline compounds. The alkalis used for this purpose include, for
14 example, alkaline aluminium salts, such as sodium aluminate, alkaline aluminium chloride
15 or alkaline aluminium nitrate, or bases, such as sodium hydroxide solution or ammonia, or
16 a combination of these alkalis.
- 17 The total quantity of aluminium added to the suspension by way of the various aluminium
18 compounds is 2.0 to 7.5% by weight, preferably 3.5 to 7.5% by weight, calculated as A_2O_3 ,
19 referred to TiO_2 base material.
- 20 0.1 to 1% by weight, preferably 0.2 to 0.5% by weight, of a magnesium compound is then
21 added, calculated as MgO and referred to TiO_2 base material. Suitable for use as the
22 magnesium compound are water-soluble magnesium salts, such as magnesium sulphate,
23 magnesium chloride and other magnesium salts, as well as mixtures of these compounds.
24 The pH value should be maintained at 8 to 10, preferably at 8.5 to 9.5, and most preferably
25 approximately 8 with the help of appropriate alkaline media, if necessary.

1 The post-treated TiO₂ pigment is subsequently separated from the suspension by filtration,
2 and the resultant filter cake is washed.

.3 To further improve the greying resistance, the pigment can additionally be treated with
4 nitrate at a concentration of up to 1.0% by weight in the finished pigment.

5 Moreover, the final pH value of the pigment may be set by adding a suitable pH modifying
6 substance such as an acid, a base, an acid salt, or a basic salt, or a combination of
7 suitable substances. The pH value is controlled by means of the degree of acidity and the
8 added quantity of the substance. In principle, all compounds may be used which do not
9 impair the optical pigment properties, which are temperature resistant during the final
10 pigment drying or pigment milling and which can be added to the filter paste, into the dryer
11 or during steam jet milling. For instance, acids like sulphuric acid, nitric acid, hydrochloric
12 acid or citric acid or acidic salts like chlorides, sulfates or the like are suitable if they comply
13 to the conditions specified.

14 Nitrate compounds are particularly suitable. By utilizing sodium nitrate the final pH value
15 comes to more than 9. Yet, the decrease of the pH value can be achieved by the use of
16 acidic nitrate compounds or a combination of acidic and non-acidic nitrate compounds as
17 for instance aluminium nitrate, a combination of aluminium nitrate and sodium nitrate, a
18 combination of aluminium nitrate and nitric acid and so forth. For example, the addition of
19 aluminium nitrate in a quantity of 0.4% by weight calculated as NO₃ in the filter paste
20 results in a lowered final pH value of from 7.5 to 8.5, preferably approximately 8.

21 Finally, the pigment is dried and milled.

22 In comparison with the reference pigments, the pigment produced according to this method
23 displays improved hiding power and improved greying resistance in the laminate, as well
24 as good retention, and is outstandingly suitable for use in decorative laminating paper.

1 Examples:

2 An example of the invention is described below. Unless otherwise stated, the quantity data
3 refer to TiO₂ base material in the suspension.

4 Example 1

5 After sand-milling, a suspension of titanium dioxide from the chloride process with a TiO₂
6 concentration of 400 g/l is adjusted to a pH value of 10 with NaOH at 60 °C. 2.4% by
7 weight P₂O₅ in the form of disodium hydrogenphosphate solution is added to the
8 suspension while stirring. The solution is added over a period of 60 minutes. After further
9 stirring for 30 minutes, 0.2% by weight TiO₂ in the form of titanyl sulphate solution is then
10 added. This is followed by further stirring for 30 minutes. In the next step, 2.7% by weight
11 Al₂O₃ in the form of acidic aluminium sulphate solution is mixed into the suspension within
12 30 minutes. After stirring for 30 minutes, the acidic suspension is set to a pH value of 9.0
13 with the help of an alkaline sodium aluminate solution in a quantity of 3.7% by weight,
14 calculated as Al₂O₃. The solution is added over a period of 40 minutes. After stirring for 30
15 minutes, 0.5% by weight MgO in the form of a magnesium sulphate solution is added. After
16 further stirring for 30 minutes the suspension is set to a pH value of 9 with NaOH.

17 After being stirred for a further 2 hours, the post-treated TiO₂ suspension is filtered and
18 washed. Following the addition of 0.25% by weight NO₃ in the form of NaNO₃, referred to
19 TiO₂ pigment, the washed filter paste is dried in a spray drier and subsequently steam-
20 milled.

21 Comparative example 1

22 The pigment is produced in a manner comparable to that described in Example 1, except
23 that titanyl sulphate and magnesium sulphate are not components of the post-treatment.
24 While stirring, 2.4% by weight P₂O₅ in the form of disodium hydrogenphosphate solution is
25 added to the sand-milled TiO₂ suspension (400 g/l TiO₂), which has a temperature of 60 °C
26 and a pH value of 10. In the next step, 3.0% by weight Al₂O₃ is mixed into the suspension

1 in the form of acidic aluminium sulphate solution. The acidic suspension is set to a pH
2 value of 7.2 with the help of an alkaline sodium aluminate solution in a quantity of 3.4% by
3 weight, calculated as Al_2O_3 . The further processing steps (filtration, washing, nitrate
4 treatment, drying, milling) are the same as in Example 1.

5 Comparative example 2

6 The pigment is produced in a manner comparable to that described in Example 1, except
7 that titanyl sulphate is not a component of the post-treatment.

8 While stirring, 2.4% by weight P_2O_5 in the form of disodium hydrogenphosphate solution
9 is added to the sand-milled TiO_2 suspension (400 g/l TiO_2), which has a temperature of
10 60 °C and a pH value of 10. In the next step, 2.6% by weight Al_2O_3 is mixed into the
11 suspension in the form of acidic aluminium sulphate solution. The acidic suspension is set
12 to a pH value of 9.2 with the help of an alkaline sodium aluminate solution in a quantity of
13 3.0% by weight, calculated as Al_2O_3 . This is followed by the addition of 0.5% by weight
14 MgO in the form of magnesium sulphate solution. NaOH is used to set a pH value of 9. The
15 further processing steps are the same as in Example 1 and Comparative example 1.

16 Test methods

17 The titanium dioxide pigments produced as described above were incorporated into
18 decorative laminating paper based on melamine resin and subsequently tested with regard
19 to their optical properties and greying resistance in pressed laminates. To this end, the
20 titanium dioxide pigment to be tested was incorporated into cellulose, and sheets with a
21 sheet weight of roughly 100 g/m² and a TiO_2 content of about 40% by mass were
22 produced.

23 a) Laminate production (laboratory scale)

24 A 36.5% aqueous pigment suspension made of 146 g titanium dioxide pigment and 254
25 g tap water is prepared. Testing is based on 30 g pulp (oven-dry). The corresponding
26 quantity of pigment suspension is adapted to the retention and the required ash content,

1 40 ± 1% in this case, and the grammage, 100 ± 1 g/m² in this case. A person skilled in the
2 art is familiar with the procedure and the auxiliaries used.

3 The ash content (titanium dioxide content) of a sheet and the retention of the pigment are
4 subsequently determined. The ash content is determined by incinerating a defined weight
5 of the produced paper in a rapid incinerator at 900 °C. The TiO₂ content by mass
6 (equivalent to the ash content) can be calculated by weighing the residue.

7 The retention is defined as the capacity to retain all inorganic substances in the sheet of
8 paper on the wire screen of the paper-making machine. The "one-pass retention" indicates
9 the percentage retained during a single feeding step to the paper-making machine. The
10 ash content in percent referred to the percentage by mass of the pigment used relative to
11 the total solids in the suspension yields the retention.

12 The further processing of the paper encompasses its impregnation and pressing into
13 laminates. The sheet to be impregnated with resin is immersed in a resin solution and pre-
14 condensed for 25 seconds at 130 °C in a recirculating-air drying oven. Impregnation is
15 performed a second time in similar manner, where the dwell time in the drying oven is 110
16 seconds. The sheet has a residual moisture content of 4 to 6% by weight. The condensed
17 sheets are combined into stacks with phenolic resin-impregnated core papers, and white
18 and black underlay paper.

19 The laminate structure used for the test comprised 9 layers: décor sheet, décor sheet, core
20 paper, core paper, black underlay, core paper, core paper, black/white underlay, décor
21 sheet.

22 The stacks are pressed for 300 seconds with the help of a Wickert Type 2742 laminating
23 press at a temperature of 140 °C and a pressure of 90 bar.

24 b) Testing

1 The optical properties and the greying resistance of the laminates were measured using
2 commercially available equipment (spectrophotometer, Xenotest weathering machine).

3 In order to assess the optical properties of laminates, the optical values (CIELAB L*, a*,
4 b*) to DIN 6174 are measured with the help of the ELREPHO® 3000 colorimeter over
5 white and black underlay. The opacity is a measure of the light transmission of the paper.

6 The following parameters were selected as a measure of the opacity of the laminates:
7 CIELAB L*_{black}, the brightness of the laminates measured over black underlay paper, and
8 the opacity value L [%] = $Y_{\text{black}}/Y_{\text{white}} \times 100$, determined from the Y-value measured over
9 black underlay paper (Y_{black}) and the Y-value measured over white underlay paper (Y_{white}).

10 The values are measured using a spectrophotometer (ELREPHO® 3000).

11 To assess the greying resistance (light-fastness) of the titanium dioxide pigments or
12 titanium dioxide pigment blends, the corresponding laminate samples are exposed in a
13 XENOTEST® 150S. The side of the laminate on which two papers are laminated together
14 is measured for the assessment. The CIELAB L*, a* and b* optical values to DIN 6174 are
15 measured before and after 96 hours of exposure in the XENOTEST® 150S. The light
16 source is a xenon-arc lamp. The temperature inside the device is 23 ± 3 °C, the relative
17 humidity being 65 ± 5%. The samples are rotated during the exposure cycle. Both $\Delta L^* =$
18 $L^*_{\text{before}} - L^*_{\text{after}}$ and $\Delta E^* = ((\Delta L^*)^2 + (\Delta a^*)^2 + (\Delta b^*)^2)^{1/2}$ are stated as a measure of the greying
19 resistance.

20 Test results:

21 The test results for the laminates produced using the pigment according to the invention
22 from the Example 1 and the pigments from Comparative examples 1 and 2 are
23 summarised in Table I. All three examples and comparative examples are set to the same
24 ash content.

25 It can be seen that the laminate produced using the pigment according to the invention
26 (Example 1) is characterised by both high opacity (L^*_{black} and L) and high greying resistance

1 (ΔL* and ΔE*). In contrast, the laminates produced using the two reference pigments 1 and
2 display significantly lower values for either opacity (Comparative example 1) or greying
3 resistance (Comparative example 2). In addition, the retention of the paper produced using
4 the pigment according to the invention was improved relative to Comparative example 2.

5 Obviously, many modifications and variations of the present invention are possible in light
6 of the above teachings. It is therefore to be understood that, within the scope of the
7 appended claims, the invention may be practiced otherwise than as specifically described.